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RESEARCH ARTICLE

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The GH5 1,4- β -mannanase from *Bifidobacterium animalis* subsp. *lactis* BI-04 possesses a low-affinity mannan-binding module and highlights the diversity of mannanolytic enzymes

Johan Morrill¹, Evelina Kulcinskaja¹, Anna Maria Sulewska^{2,4}, Sampo Lahtinen³, Henrik Stålbrand¹, Birte Svensson² and Maher Abou Hachem^{2*}

Abstract

Background: β -Mannans are abundant and diverse plant structural and storage polysaccharides. Certain human gut microbiota members including health-promoting *Bifidobacterium* spp. catabolize dietary mannans. Little insight is available on the enzymology of mannan deconstruction in the gut ecological niche. Here, we report the biochemical properties of the first family 5 subfamily 8 glycoside hydrolase (GH5_8) mannanase from the probiotic bacterium *Bifidobacterium animalis* subsp. *lactis* BI-04 (BMan5_8).

Results: BMan5_8 possesses a novel low affinity carbohydrate binding module (CBM) specific for soluble mannan and displays the highest catalytic efficiency reported to date for a GH5 mannanase owing to a very high k_{cat} ($1828 \pm 87 \text{ s}^{-1}$) and a low K_m ($1.58 \pm 0.23 \text{ g} \cdot \text{L}^{-1}$) using locust bean galactomannan as substrate. The novel CBM of BMan5_8 mediates increased binding to soluble mannan based on affinity electrophoresis. Surface plasmon resonance analysis confirmed the binding of the CBM10 to manno-oligosaccharides, albeit with slightly lower affinity than the catalytic module of the enzyme. This is the first example of a low-affinity mannan-specific CBM, which forms a subfamily of CBM10 together with close homologs present only in mannanases. Members of this new subfamily lack an aromatic residue mediating binding to insoluble cellulose in canonical CBM10 members consistent with the observed low mannan affinity.

Conclusion: BMan5_8 is evolved for efficient deconstruction of soluble mannans, which is reflected by an exceptionally low K_m and the presence of an atypical low affinity CBM, which increases binding to specifically to soluble mannan while causing minimal decrease in catalytic efficiency as opposed to enzymes with canonical mannan binding modules. These features highlight fine tuning of catalytic and binding properties to support specialization towards a preferred substrate, which is likely to confer an advantage in the adaptation to competitive ecological niches.

Keywords: *Bifidobacterium*, Carbohydrate-binding module, Gut microbiota, Mannan, Probiotic bacteria, Surface plasmon resonance

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Background

β -Mannans (hereafter mannans) are abundant polysaccharides playing diverse roles in plants including energy storage in seed endosperms, e.g. carob and guar seeds, legumes, coconuts and coffee beans [1], or structural support in the hemicellulose cell wall matrix [2], where they can constitute up to 25 % of the dry mass in softwood. The biochemical details of enzymatic mannan depolymerization have received increasing attention in recent years due to wide interest within the biofuel and biorefinery areas [3, 4]. Mannans, however, are also present in human nutrition, either as cell wall components in cereal grains and some fruits and vegetables such as kiwi, apple and tomato [5–8] or as common hydrocolloid food additives used as thickeners or to adjust texture [9, 10]. Mannans are not known to be hydrolyzed by human digestive enzymes and thus offer a potential resource to mannanolytic gut bacteria. The fermentation of guar gum (GG) galactomannan has been demonstrated in the human gut [11] and intake of partial hydrolysates of this polysaccharide stimulated the proliferation of *Bifidobacterium* spp. in humans [12] and mice [13]. However, insight into the microbial strategies and enzymes mediating mannan degradation in the human gut lags behind.

Mannans occur as insoluble crystalline polymers of β -1,4-linked mannosyl residues in seeds, such as ivory nut mannan (INM) or as soluble heteropolymeric glucomannans consisting of alternating β -1,4-linked glucosyl and mannosyl backbone e.g. in corms of *Amorphophallus konjac*. Mannan can be substituted to different degrees with α -1,6-linked galactosyl side-chains as in carob (locust bean) and guar gums and in glucomannan from softwood, which also has acetyl decorations [14]. Konjac glucomannan (KGM), locust bean (LBG) and GG galactomannans are the main mannans used as food additives [9, 10].

The concerted action of several backbone- and sidechain-degrading enzymes is required for depolymerization of mannans. The precise number of enzymatic activities varies with the substrate structure, but endo- β -1,4-mannanases (EC 3.2.1.78) that hydrolyze internal backbone β -1,4-linkages are central in mannan degradation. β -Mannanases are assigned in glycoside hydrolase (GH) families 5, 26 and 113 in the Carbohydrate-Active enZymes (CAZy) database [15]. Many bacterial mannanases cluster in subfamily 8 of GH5 (GH5_8) according to a phylogeny-based assignment [16]. Mannanases of GH5 employ a double displacement mechanism with the retention of anomeric configuration [17]. Some mannanases contain carbohydrate-binding modules (CBMs) that have been ascribed diverse roles including targeting enzymes to polysaccharides, increasing the local substrate concentration, or conferring processivity [18, 19].

The present study focuses on the widely utilized and clinically well-documented probiotic bacterium *Bifidobacterium animalis* subsp. *lactis* Bl-04 [20]. Genome analysis identified a gene encoding a mannanase comprising a GH5 catalytic module joined to a CBM10. Members of CBM10 were previously found to bind to insoluble microcrystalline cellulose [21] and insoluble mannan [22]. Furthermore, we show that the GH5 mannanase (BMan5_8) from *Bifidobacterium animalis* subsp. *lactis* Bl-04, which is conserved within *Bifidobacterium animalis* subsp. *lactis*, displays the highest catalytic efficiency reported to date for a GH5 β -mannanase owing to a combination of very high k_{cat} and low K_m . The CBM10 of this enzyme is the first described low-affinity mannan binding module and it forms a novel CBM10 subfamily together with close homologues. The distinct differences in the biochemical properties of this enzyme as compared to characterized β -mannanases from gut microbiota illustrate the diversity of mannan utilization strategies, which is likely to be important in adaptation to the highly competitive gut niche.

Methods

Carbohydrates

Cellotetraose, locust bean gum (LBG), microcrystalline cellulose (Avicel) and hydroxyethyl cellulose (HEC) are from Sigma-Aldrich (St. Louis, MO, USA); mannotriose (M_3), mannotetraose (M_4), mannopentaose (M_5), mannohexaose (M_6), low-viscosity locust bean gum (LBG-lv), ivory nut mannan (INM) and konjac glucomannan (KGM) are from Megazyme (Bray, Ireland); guar gum (GG) is from Carl Roth (Karlsruhe, Germany). The compositions of the polysaccharides are listed in Additional file 1.

Cloning

Bifidobacterium animalis subsp. *lactis* Bl-04 chromosomal DNA [23] was used to clone the locus Balac_1450 (GenBank accession number ACS46797) encoding a β -mannanase of glycoside hydrolase family 5 subfamily 8 (GH5_8). The PCR-amplified gene fragment encoding the full-length mature β -mannanase lacking the N-terminal signal peptide (amino acid residues 1–29 as predicted by Signal P v.4.0, <http://www.cbs.dtu.dk/services/SignalP/>), referred to as BMan5_8, was cloned within the *NheI* and *BamHI* restriction sites in pET28a(+) (Novagen, Darmstadt, Germany) using a primer pair (Additional file 2) to yield the plasmid pET28a-BMan5_8. This plasmid, encoding the enzyme fused to an N-terminal hexa-histidine purification tag, was transformed first into *Escherichia coli* TOP10 (Life Technologies, Grand Island, NY, USA). The presence of the insert was verified by sequencing and restriction analysis. The construct encoding the

truncated enzyme *B/Man5_8-ΔCBM10* lacking the C-terminal CBM10, was generated by site-directed mutation of Ser³³⁸, which is located approximately in the middle of the linker sequence (G³³⁰GGNSGGS GNTGGNSGTTDDG³⁵¹) between the catalytic and the binding modules, into a stop codon (QuikChange mutagenesis kit; Agilent Technologies, Santa Clara, CA, USA) and the primer pair in Additional file 2 to yield pET28a-*B/Man5_8-ΔCBM10*. The mutation was verified by full sequencing. The plasmids encoding *B/Man5_8* and *B/Man5_8-ΔCBM10* were transformed into *E. coli* BL21 (DE3) for production.

Enzyme production and purification

Production of *B/Man5_8* and *B/Man5_8-ΔCBM10* was carried out in 2 L baffled shake flasks. Overnight cultures were used to inoculate 1 L LB medium containing 10 mM glucose and 50 μg·mL⁻¹ kanamycin and grown at 30 °C to an OD₆₀₀ of 0.5. Protein expression was induced with 100 μM IPTG and growth was continued for 5 h. For *B/Man5_8*, the cells were harvested by centrifugation, resuspended in binding buffer (10 mM HEPES, 0.5 M NaCl, 10 % glycerol, 15 mM imidazole, pH 7.4) with added protease inhibitor cocktail (Roche Diagnostics, Indianapolis, IN) and lysed using a high pressure homogenizer, centrifuged and filtered through a 0.45-μm filter. For *B/Man5_8-ΔCBM10*, the cells were lysed using the BugBuster™ Protein Extraction Reagent (Novagen), according to the manufacturer's recommendations. Clarified *B/Man5_8* and *B/Man5_8-ΔCBM10* lysates were applied onto a 5-mL HisTrap HP (GE Healthcare, Uppsala, Sweden), washed with binding buffer, and eluted with a gradient formed with 400 mM imidazole in 10 mM HEPES, 0.5 M NaCl, 10 % glycerol, pH 7.4. Eluted fractions were analyzed by SDS-PAGE and for β-mannanase activity as described below. Pure fractions were pooled and concentrated (10 kDa cut-off ultrafiltration units; Amicon), and the histidine tag was cleaved off using human plasma thrombin (Calbiochem, San Diego, CA, USA) according to the manufacturer's instructions. Cleaved enzymes were recovered after passing through a binding buffer pre-equilibrated 1 mL HisTrap HP column (GE Healthcare). The concentrated enzyme samples were further purified to electrophoretic homogeneity by chromatography on a HiLoad Superdex G75 26/60 column (GE Healthcare) eluted with 10 mM HEPES, pH 7.0 in 1.2 column volumes. The pure enzyme samples were concentrated as above and stored at 4 °C until further use.

Basic enzymatic properties and enzyme stability

β-Mannanase activity was measured towards 2.5 g·L⁻¹ LBG at 37 °C for 10 min in 400 μL standard assay buffer (40 mM phosphate-citrate buffer, pH 6.0, with 0.005 %

(v/v) Triton X-100) with appropriately diluted enzyme. The substrate preparation and reducing end 3,5-dinitrosalicylic acid (DNS) assay were performed as previously reported [24, 25]. The pH dependence of activity was initially determined using 6.4 nM *B/Man5_8* with LBG in 100 mM Britton-Robinson buffers at pH 2–10 and then refined in 40 mM phosphate-citrate at pH 5–7. To assess storage stability as function of pH, 1.28 μM *B/Man5_8* was incubated at 4 °C for four days in 100 mM Britton-Robinson buffers at pH 2–10, thereafter diluted to 3.2 nM in standard assay buffer, and residual β-mannanase activity was determined as described above. The dependence of enzyme activity on temperature was determined for 6.4 nM *B/Man5_8* with LBG in standard assay buffer, in the range 25–80 °C.

The conformational stability of 12.6 μM *B/Man5_8* in 10 mM MES buffer, pH 6.5 was assessed using differential scanning calorimetry (DSC) between 15 °C and 90 °C at 1 °C·min⁻¹ in a MicroCal VP-DSC calorimeter. The enzyme was scanned twice with cooling and pre-equilibration in between to assess the reversibility of thermal transitions. The dialysis buffer was used as a reference and the reference and baseline corrected thermograms were analyzed using Origin7 with a DSC add-on provided with the instrument.

Substrate specificity

The soluble substrates LBG, LBG-lv, GG and KGM were prepared as previously described [25]. The insoluble substrates INM and Avicel were washed three times in water, twice for 2 h and once overnight, with a 15:1 volume to mass ratio, followed by 3 h wash in buffer. The specific activity of *B/Man5_8* towards LBG-lv and KGM (2.5 g·L⁻¹, 10 min incubation), as well as GG (2.5 g·L⁻¹, 2.5 h incubation), INM (5 g·L⁻¹, 45 min incubation) and Avicel (10 g·L⁻¹, 3 h incubation) was measured using the standard activity assay.

Enzyme kinetics

LBG-lv (0.45–9.0 g·L⁻¹) was hydrolyzed by 1 nM *B/Man5_8* or *B/Man5_8-ΔCBM10* at 37 °C in standard assay buffer as measured by DNS (see above). The initial reaction rates were determined and the kinetic parameters k_{cat} and K_m were determined by fitting the Michaelis-Menten equation to the data using Prism 6 (GraphPad Software, La Jolla, CA, USA). Experiments were performed in duplicates.

For kinetic analysis of M₅ hydrolysis, 0.25 nM *B/Man5_8* was incubated with 0.25–10 mM M₅ at 37 °C in standard assay buffer. Samples were withdrawn at 0, 5, 10, 15 and 20 min, and boiled for 5 min. The hydrolysis products of M₅ were analyzed by HPAEC-PAD using a DX-500 system (Thermo Scientific Dionex, Sunnyvale, CA, USA). A CarboPac™ 100 column and guard column

were used with a mobile phase of 78 mM sodium hydroxide at a flow rate of $1.0 \text{ mL} \cdot \text{min}^{-1}$. The decrease in M_5 concentration is stoichiometrically equivalent to the increase in concentrations of degradation products. These concentrations were calculated and used to determine k_{cat} and K_m as described [26].

Mannan hydrolysis action patterns

The product profiles for 5.3 nM *B/Man5_8* or *B/Man5_8-ΔCBM10* towards $2.5 \text{ g} \cdot \text{L}^{-1}$ INM or LBG in standard assay buffer at 37 °C at 0 min, 30 s, 5, 15 and 30 min, as well as 2 and 24 h, were analyzed by HPAEC-PAD (ICS-5000 system; Thermo Scientific Dionex, Sunnyvale, CA, USA). A CarboPac™ PA-200 column and guard column were used with a mobile phase of 78 mM sodium hydroxide and a linear gradient of 0–50 mM sodium acetate at a flow rate of $0.5 \text{ mL} \cdot \text{min}^{-1}$.

Binding to polysaccharides and oligosaccharides

Qualitative screening of binding of *B/Man5_8* and *B/Man5_8-ΔCBM10* to soluble polysaccharides was analysed using affinity electrophoresis as previously described [27]. The electrophoretic separation of 2 µg of either protein was performed at 4 °C and 45 V for 20 h in 50 mM Tris-Borate, pH 8.7 and in 12 % polyacrylamide gels without (control) or with 0.1, 0.5 or $2.5 \text{ g} \cdot \text{L}^{-1}$ KGM, as well as $2.5 \text{ g} \cdot \text{L}^{-1}$ hydroxyethyl cellulose (HEC). Quantitative binding analysis of *B/Man5_8* and *B/Man5_8-ΔCBM10* to LBG-lv was done using affinity electrophoresis [27, 28] at 100 V for 4 h in gels without (control) or with 0.05, 0.10 or $0.25 \text{ g} \cdot \text{L}^{-1}$ LBG-lv, as described above. The apparent dissociation constant (K_d) for LBG-lv was calculated by nonlinear regression using Prism 6 (GraphPad Software, La Jolla, CA, USA). Binding to insoluble INM and Avicel was measured by adding 50 µl of appropriately diluted enzyme to 400 µl of 0 (negative control), 0.5, 5, 50 or $100 \text{ g} \cdot \text{L}^{-1}$ of either substrate suspended in standard assay buffer. The slurry was incubated at 4 °C for 1 h with gentle shaking, centrifuged at $16\,000 \times g$ for 10 min and the mannanase activity in the supernatant was assayed as described above.

Surface plasmon resonance (SPR) analysis was performed using a Biacore T100 (GE Healthcare) to assess binding to manno-oligosaccharides and cellotetraose. Enzymes diluted to $40 \text{ µg} \cdot \text{mL}^{-1}$ in 10 mM sodium acetate, pH 3.9, were immobilized onto a CM5 chip (GE Healthcare) using a standard amine coupling protocol to a density of 3999 and 2879 response units (RU) for *B/Man5_8* or *B/Man5_8-ΔCBM10*, respectively. Binding analysis was performed in triplicates at 25 °C in 10 mM MES, 150 mM NaCl, pH 6.0, 0.005 % (v/v) P20 (GE Healthcare) running buffer at $30 \text{ µL} \cdot \text{min}^{-1}$ with association and dissociation times of 60 and 30 s, respectively, at

seven concentrations of M_4 (0.5–25 mM), M_5 (0.05–25 mM), M_6 (0.05–12.5 mM) and cellotetraose (0.05–25 mM). A one-binding-site model was fit to the blank and reference cell-corrected steady state response data to determine the dissociation constant K_d and the saturation binding level R_{max} , which was used to estimate the molar stoichiometry (n) [29] of oligosaccharide binding to immobilized enzymes that retain binding activity.

Action pattern analysis using mannopentaose

For analysis of the productive binding modes of M_5 , enzymatic reactions were carried out in ^{18}O labeled water (H_2^{18}O 97 % pure from Sigma-Aldrich, St. Louis, MO, USA), as described [26, 30] at 8 °C with 1 mM M_5 and 20 nM *B/Man5_8* in 1 mM sodium phosphate, pH 6.0 in 93 % H_2^{18}O . Samples were incubated for 0–24 h and analyzed with MALDI-TOF MS. Aliquots of 0.5 µL were withdrawn at each time point, spotted directly on a MALDI plate, covered with 0.5 µL matrix (10 g $\cdot \text{L}^{-1}$ 2,5-dihydroxybenzoic acid (DHB) in water) and dried under warm air. This analysis combined with HPAEC-PAD quantification of M_3 and M_4 products of M_5 hydrolysis was used to determine the frequency of productive binding modes of M_5 as previously described [26].

Transglycosylation product formation from M_5

For analysis of transglycosylation 5 mM M_5 was incubated with 1.6 nM *B/Man5_8* in standard assay buffer for 0–24 h. Samples were analyzed with MALDI-TOF MS as described above.

Sequence analyses and homology modeling

The sequence of *B/Man5_8* was analyzed to identify catalytic and ancillary modules using dbCAN [31], a CAZy-based database [15] for automated carbohydrate-active enzyme annotation. The sequence of the *B/Man5_8* catalytic module was aligned with 69 sequences listed as GH5_8 in CAZy [15], as well as 12 characterized β-mannanases from GH5 subfamilies 7, 8 and 10, using MAFFT 7 [32]. The CBM10 of *B/Man5_8* was aligned with 83 predicted CBM10 sequences retrieved using dbCAN, after removing two sequences that appeared fragmented and three sequences that displayed E-values above 0.02. These sequences were aligned using MAFFT 7 and a phylogenetic tree was calculated using the unweighted pair group method with arithmetic mean using standard settings accessed on the CBRC server (<http://mafft.cbrc.jp/alignment/server/phylogeny.html>) and rendered using Dendroscope 3 [33]. The crystal structure of the *Thermobifida fusca* mannanase TjMan5A (PDB ID: 2MAN) [34] served as template to model the 39 % amino acid sequence identical catalytic module of *B/Man5_8*, using MODELLER [35]. The homology model was evaluated with MolProbity [36], showing that 93.29 %

of all residues were in Ramachandran favored regions. The model had a clash score of 72.89, defined as the number of unfavorable (≥ 0.4 Å) steric overlaps per 1000 atoms [37]. Evaluation with ProQ [38] gave a predicted LGscore of 5.903 and a predicted MaxSub of 0.266, indicating good model quality.

Results

Basic properties of *B/Man5_8*

B/Man5_8 is predicted to be an extracellular modular β -mannanase comprising a catalytic module of glycoside hydrolase family 5 subfamily 8 (GH5_8) joined to a C-terminal carbohydrate-binding module of family 10 (CBM10) by a typical linker sequence plausibly starting from Gly³³⁰ to Gly³⁵¹ (see sequence in Methods). The position of the truncation (Ser³³⁸) was chosen as to avoid destabilisation of the catalytic module due to possible loss of interactions with the first few residues in the linker. The full-length *B/Man5A* and the truncated *B/Man5_8*- Δ CBM10 were produced and purified to electrophoretic homogeneity. Both enzyme forms migrated in accord with their theoretical molecular mass of 37.9 kDa and 33.8 kDa, respectively. The lack of activity towards microcrystalline cellulose (Avicel) and the high activity towards mannans (Table 1) confirmed the β -mannanase specificity of *B/Man5_8*.

B/Man5_8 has highest activity at pH 6.0 and retains approximately 75 % of its maximal activity between pH 5 and 7 using LBG as a substrate (Additional file 3A). After incubation for 4 days at 4 °C, the enzyme retained 100 % of its initial activity at pH 8.0, and approximately 75 % at pH 6.0 (Additional file 3B). *B/Man5_8* displayed highest β -mannanase activity at 55 °C and pH 6.0 (Additional file 4A), corresponding to an Arrhenius activation energy of 69.5 kJ·mol⁻¹ (Additional file 4B). The unfolding temperature of *B/Man5_8* was determined to 55.9 °C by differential scanning calorimetry (DSC) analysis asserting the structural integrity of the enzyme. The unfolding thermogram featured a single transition (Additional file 5), suggesting that the unfolding of the two modules of *B/Man5_8* is overlapping.

Table 1 Specific activity of *B/Man5_8* towards soluble and insoluble polysaccharides

Substrate	Specific activity (kat·mol ⁻¹)	Specific activity (U·mg ⁻¹)
LBG	1213	1920
LBG-lv	872	1380
GG	n.d. ^a	n.d. ^a
KGM	1592	2520
INM	76	120
Avicel	n.d. ^a	n.d. ^a

^a Not detected

Substrate specificity and kinetics

Activity of *B/Man5_8* was highest towards the soluble KGM followed by LBG and low-viscosity LBG (LBG-lv) (Table 1). Activity towards insoluble INM was only 6 % of that on LBG, and no activity towards GG could be measured (Table 1). The apparent kinetic parameters for *B/Man5_8* and *B/Man5_8*- Δ CBM10 were determined for LBG-lv (Table 2; Additional file 6A) and marginal increases in K_m and k_{cat} were observed as a result of deleting the CBM10. Apparent kinetic parameters were also determined towards mannopentaose (M₅) for *B/Man5_8* (Table 2; Additional file 6B). The enzyme displayed no activity towards mannotriose (M₃), and very low activity towards mannotetraose (M₄), but was highly active on larger manno-oligosaccharides as analyzed by high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) (data not shown).

Product profiles, transglycosylation and productive binding frequency analysis

The hydrolysis profiles of LBG and INM by *B/Man5_8* and *B/Man5_8*- Δ CBM10 were analyzed using HPAEC-PAD. Both enzyme variants produced M₃, M₄ and M₅ as the main initial hydrolysis products from both LBG and INM, along with some mannobiose (M₂) (Fig. 1). The main products of *B/Man5_8* from M₅ were M₂, M₃ and M₄. The formation of 10-fold higher M₄ compared to mannose (M₁) (Fig. 2) is indicative of transglycosylation activity, which was supported by the detection of mannohexaose (M₆), mannoheptaose (M₇) and mannooctaose (M₈) transglycosylation products by mass-spectrometry following incubation with M₅ (Fig. 3). All four productive M₅ binding modes were observed, albeit with higher frequency (66 %) for binding modes where the larger part of the substrate was bound at the aglycone binding subsites (Fig. 4).

Binding to polysaccharides and oligosaccharides

Based on retardation in affinity electrophoresis, *B/Man5_8* bound to LBG-lv (Additional files 7–8) with an apparent K_d of 0.31 g·L⁻¹, while the affinity of *B/Man5_8*- Δ CBM10 was too low to be determined due to lack of change in relative retardation as a function of increasing the LBG concentration. Neither the full-length nor the truncated enzyme displayed binding to soluble hydroxyethyl cellulose (HEC), insoluble cellulose (Avicel) or insoluble mannan (INM) in this analysis.

SPR is a particularly powerful tool to quantify low affinity interactions as observed above for *B/Man5_8* binding to mannan. Ideally, CBM binding analysis would be carried out on an isolated CBM. In our case, however, the small size of the CBM10 and the presence of only a single lysine residue (used for random amine coupling

Table 2 Kinetic parameters for *B*/Man5_8 on LBG-lv and *M*₅

Substrate	Enzyme	<i>k</i> _{cat}	<i>K</i> _m	<i>k</i> _{cat} / <i>K</i> _m
LBG-lv	<i>B</i> /Man5_8	1828 ± 87 s ⁻¹	1.58 ± 0.23 g · L ⁻¹	1157 ± 177 L · s ⁻¹ · g ⁻¹
	<i>B</i> /Man5_8-ΔCBM10	2005 ± 179 s ⁻¹	1.75 ± 0.47 g · L ⁻¹	1146 ± 324 L · s ⁻¹ · g ⁻¹
<i>M</i> ₅	<i>B</i> /Man5_8	97 ± 2 s ⁻¹	1.09 ± 0.11 mM	89 ± 9 s ⁻¹ mM ⁻¹

or biotinylation), renders the immobilization of the protein to sufficient levels challenging. To circumvent this limitation and avoid using this single lysine residue for immobilization with risk of affecting the function of the module, the binding analysis was done using immobilized full-length enzyme and *B*/Man5_8-ΔCBM10 to evaluate the contribution of the CBM10 to binding. Only the full-length enzyme showed affinity towards *M*₄ based on the SPR analysis, which suggested that the CBM10 mediated binding to *M*₄ (Table 3). The apparent binding stoichiometry of both *M*₅ and *M*₆ was approximately twice for the full-length enzyme as compared to the isolated catalytic module (Table 3), reflecting binding of these ligands to both the GH5 catalytic module and the CBM10. No binding to cellotetraose was measured with either enzyme variant.

Sequence analyses and homology modeling

B/Man5_8 contains the seven amino acids that are conserved in family GH5 [39] (Additional file 9). The homology model of *B*/Man5_8 revealed that residues interacting with the substrate in subsites -4 through -2 are conserved between *B*/Man5_8 and *Tf*Man5A [34] (Fig. 5). There are also several aromatic residues in the putative aglycone binding region of *B*/Man5_8 including Trp¹⁹⁶ and Trp²²⁵ which are invariant in GH5_8 as well as Trp²⁰⁰ which is conserved in 77 % of GH5_8 sequences (Additional file 10).

The CBM10 of *B*/Man5_8 and close homologues thereof, which are exclusively joined to β-mannanase catalytic modules, populate a distinct clade in the CBM10 phylogenetic tree (Fig. 6A). This clade is distinguished by unique sequence features, including substitution of a

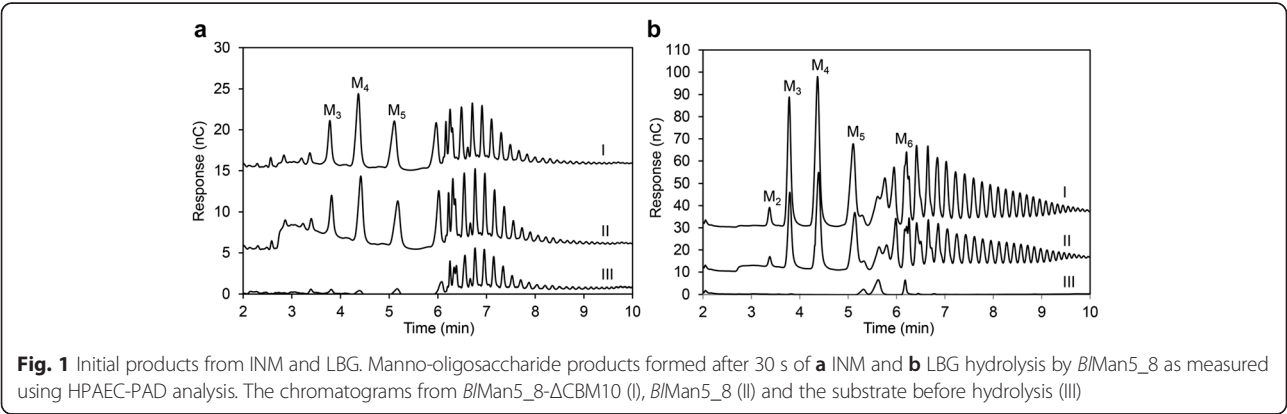
tyrosine residue conserved in other CBM10s as well as an insertion of five residues (Fig. 6B).

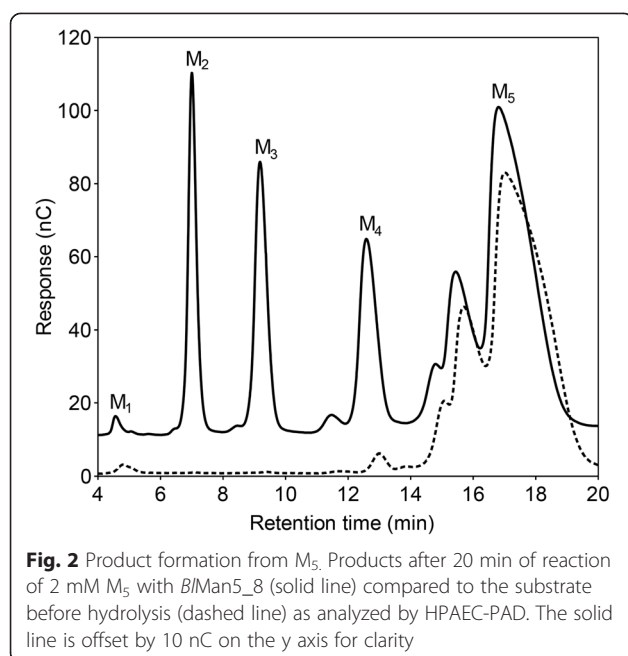
Discussion

The human gastrointestinal tract comprises one of the most densely populated ecological niches in nature, with bacterial counts exceeding 10¹¹ cells · g⁻¹ content [40]. Since most dietary glycans are indigestible to humans, their metabolism plays an instrumental role in shaping the composition of the gut microbiota [41]. To date, the GH26 mannanases from *Bacteroides fragilis* [42] and *Bifidobacterium adolescentis* [28] are the only described β-mannanases from the gut niche. In this study we report the biochemical properties of the first GH5 mannanase from the *Bifidobacterium* genus.

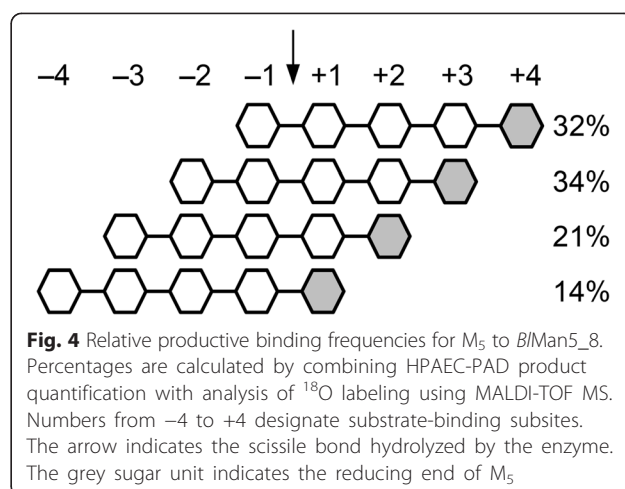
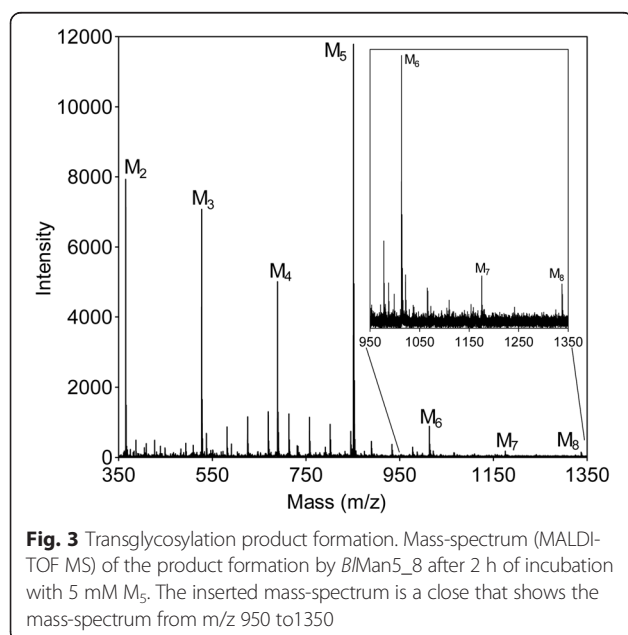
***B*/Man5_8 is an exceptionally potent enzyme optimized for efficient depolymerization of soluble mannan**

The preferred substrates for *B*/Man5_8 are soluble glucomannan and galactomannan with low degree of galactosyl substitution (Table 1). *B*/Man5_8 displays 19-fold higher *k*_{cat} on LBG as compared to *M*₅ (Table 2), suggesting that substrate contacts beyond five mannosyl residues contribute to catalytic turnover, similarly to other mannanases [43]. The catalytic efficiency of *B*/Man5_8 on LBG is the highest measured to date for a GH5 mannanase owing to both a very high *k*_{cat} and a low *K*_m (Additional file 11). Notably, the *K*_m value for *B*/Man5_8 towards soluble mannan is one of the lowest amongst bacterial mannanases, which is likely to be an important advantage in the adaptation to the highly competitive gut ecological niche. The *K*_m value in part reflects substrate affinity, which appears higher at the





aglycone-accommodating region of the active site cleft as evident from the productive binding frequency of M_5 (Fig. 3). Substrate affinity at aglycone subsites has been shown to contribute to lowering K_m and increasing transglycosylation [30, 44]. This is in line with the high transglycosylation activity of *B/Man5_8*, reflected by the disproportionate liberation of M_4 relative to M_1 ratio using M_5 as a substrate (Fig. 2), which is attributed to secondary hydrolysis of transglycosylation products of larger degree of polymerization (Fig. 3). Interestingly, sequence comparison and homology modeling revealed that the



putative aglycone binding regions of *Thermobifida fusca* GH5_8 mannanase (*T/Man5A*) [34] and *B/Man5_8* are lined with conserved aromatic residues, including Trp¹⁹⁶ and Trp²⁰⁰ (Additional file 10; Fig. 5). Indeed, the position and conservation of these residues correlate with the favored productive binding at the aglycone region (Fig. 4).

The CBM10 of *B/Man5_8* binds soluble mannan with low affinity and represents a new clade within CBM10

The CBM10 of the modular *CjXyn10A* xylanase from *Cellvibrio japonicus*, which binds tightly to Avicel, is the only structurally characterized CBM10 member [21]. The Man5A β -mannanase from the same organism, which possesses a C-terminal tandem repeat of two CBM10 modules (*CjMan5A*-CBM10-1-CBM10-2), was also shown to bind both Avicel and insoluble mannan (INM) [22]. Deletion of CBM10-2 of *CjMan5A*, which is a close homolog to the counterpart from *CjXyn10A*, resulted in a severe reduction of affinity to Avicel and INM [22], which hinted that the CBM10-1 was not efficiently mediating binding to the above insoluble substrates. Similarly, the CBM10 of *B/Man5_8*, which

Table 3 Surface plasmon resonance binding analysis of *B/Man5_8* to oligosaccharides

Enzyme	Substrate	K_d (mM)	n^a
<i>B/Man5_8</i>	M_4	5.1 ± 0.8	0.78
	M_5	3.7 ± 0.3	0.94
	M_6	0.58 ± 0.05	0.92
	Cellotetraose	n.d. ^b	–
<i>B/Man5_8</i> - Δ CBM10	M_4	n.d. ^b	–
	M_5	1.5 ± 0.3	0.36
	M_6	0.24 ± 0.42	0.53
	Cellotetraose	n.d. ^b	–

^a Estimated stoichiometry of binding to the immobilized protein

^b Not detected

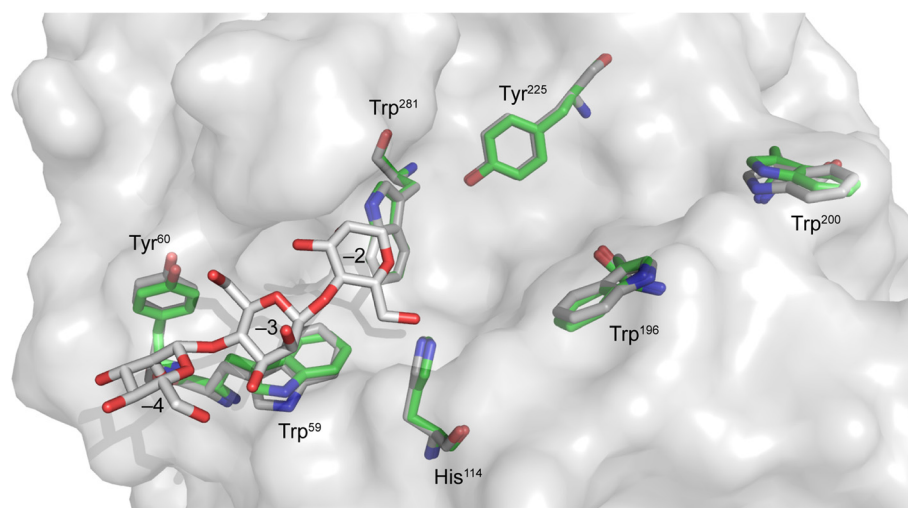


Fig. 5 Modeling of putative subsite residues in *B/Man5_8*. Conserved aromatic residues in the active site of *B/Man5_8* are visualized (green) by homology modeling using the structure from *T/Man5A* (PDB ID: 1BQC) as template (grey). A semi-transparent molecular surface is shown to depict the topology of the active site of *T/Man5A*. The conserved aromatic residues are designated with *B/Man5_8* numbering and the co-crystallized M_3 , accommodated at subsites -4 through -2 in *T/Man5A*, is shown in white

belongs to the same subgroup as the CBM10-1 of *CjMan5A*, did not bind to Avicel or HEC. By contrast, this CBM10 conferred increased affinity to soluble mannan based on larger relative retardation in affinity electrophoresis of the full-length *B/Man5_8* compared to the truncated *B/Man5_8*- Δ CBM10. Remarkably, the affinity of *B/Man5_8* for soluble galactomannan is 35 – 67-fold lower than that reported for GH26 mannanases *BaMan26A* from *Bifidobacterium adolescentis* (*BaMan26A*) [28] and *CjMan26A* from *Cellulomonas fimi* [45] possessing typical mannan-specific CBM23 modules. Binding analysis using SPR confirmed the mannan specificity and low affinity of the CBM10 of *B/Man5_8*, as deletion of this module considerably reduced the apparent stoichiometry of binding to M_5 and M_6 (Table 3). *B/Man5_8*- Δ CBM10 moreover displayed about two-fold higher affinity to M_5 and M_6 than the full-length enzyme, suggesting that the CBM10 had lower affinity than the catalytic module. The CBM10 of *B/Man5_8* is a novel low-affinity CBM that binds manno-oligosaccharides with K_d values in the mM range, in contrast to canonical moderate-affinity mannan-binding CBMs displaying K_d values in the low μ M range [46, 47]. Low-affinity starch-binding CBMs, conferring more dynamic binding, have been identified in plastidial α -amylases and starch-regulatory glucan water dikinases [48, 49].

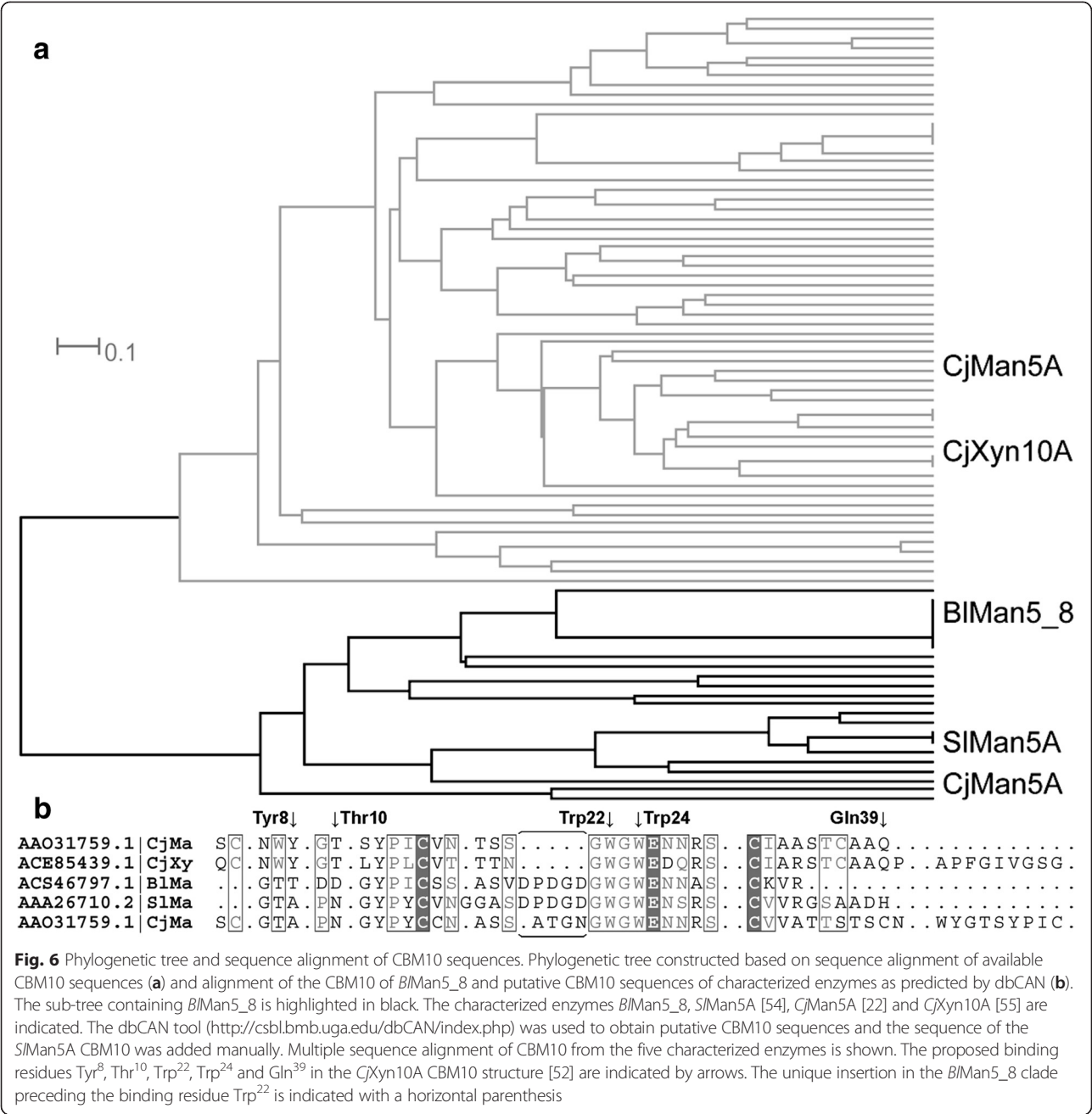
The low affinity of the CBM10 is consistent with its very modest impact on the kinetic parameters of LBG hydrolysis (Table 2). The trend of increased K_m and k_{cat} upon CBM deletion is similar to other enzymes acting on soluble substrates, but the magnitude of these

changes is much larger with typical CBMs than for the CBM10 of *B/Man5_8* [50, 51].

The CBM10 of *B/Man5_8* clusters exclusively with counterparts from GH5 and GH26 mannanases and segregates in the phylogenetic tree from sequences resembling the cellulose-specific CBM10 from *CjXyn10A* (Fig. 6; Additional file 12) also occurring in xylanases, cellulases and other carbohydrate-active enzymes as well as mannanases (Additional file 12). The aromatic binding residue Tyr⁸ in the cellulose-binding site of CBM10 of *CjXyn10A* [21] is substituted to smaller residues (Fig. 6, Additional file 9) in the *B/Man5_8* CBM10 subfamily, that possesses a unique insertion preceding the substrate binding residue Trp²² and lacks the two conserved cysteines Cys⁵ and Cys³⁸ (Fig. 6) suggesting the loss of one disulphide bridge compared to the clade represented by the CBM10 of *CjXyn10A* [52] (Additional file 13). Altogether, the loss of the Tyr⁸ aromatic stacking platform combined with significant differences in the loop length close to the binding Trp²² provide a possible explanation for the low affinity of the CBM10 from *B/Man5_8* compared to *CjXyn10A*. Additionally, the loss of a disulfide bridge may elicit flexibility and/or structural changes that affect the biochemical properties of the *B/Man5_8* CBM10 and its close homologs. Assigning the effects of these differences requires further structural and mutational analyses of this clade of CBM10.

Mannan utilization in the gut and the biological implication of *B/Man5_8* activity

The concentration and residence time of mannan in the gastrointestinal tract is expected to be highly dynamic



and dependent on nutritional intake and gut microbiota composition. The presence of mannan utilization pathways in human gut-adapted taxa, including the major gut commensal genus *Bacteroides*, reflects an evolutionary adaptation to the dietary intake of this glycan. Recently, a novel pathway for the utilization of mannan involving a putative extracellular mannobiose-forming GH26 exomannanase from *Bacteroides fragilis* (*BfMan26A*) was reported [42]. The mannobiose released by this enzyme is taken up by specialized ion symporters and subsequently degraded and metabolized intracellularly by an epimerase

and a phosphorylase. This GH26 enzyme does not possess a CBM, and was speculated to act on mannan fragments produced by other β -mannanases from the same organism. The extracellular putatively cell-attached GH26 mannanase from *Bifidobacterium adolescentis* (*BaMan26A*) hydrolyzes mannan to mainly mannotriose [28]. This modular enzyme, featuring a GH26 catalytic module joined to a C-terminal tandem repeat of CBM23 mannan-binding modules, binds tightly to LBG ($K_d = 8.8 \text{ mg} \cdot \text{L}^{-1}$) [28]. The deletion of the tandem CBM23 repeat abolishes measurable binding and yields an enzyme with very high

K_m ($21.3 \text{ g} \cdot \text{L}^{-1}$), reminiscent of typical canonical moderate-affinity CBMs that decrease apparent K_m values of the enzymes they are attached to [50, 51]. This CBM-mediated decrease in K_m , however, is often associated with a penalty of reduced k_{cat} [50, 51]. Such effects are very different in case of *B/Man5_8*, where the CBM10 makes little contribution to the low K_m . The justification for maintaining a CBM, albeit with lower affinity, could be a trade-off to increase the substrate-binding affinity of the enzyme (as observed from retardation electrophoresis data) and thus lower the k_{off} of the polymeric substrate, while minimizing the energy barrier pertaining to anchoring of the substrate tightly to the CBM. This may maintain proximity of the substrate to the enzyme, which has been shown to promote additional hydrolysis events following the initial enzyme-substrate encounter in cellulases [53] due to a higher $k_{\text{cat}}/k_{\text{off}}$ ratio. The optimization of substrate affinity through a low-affinity CBM is likely to offer a different adaptation solution to the fierce competition prevalent in the gut niche as opposed to the higher substrate affinity, but decrease in catalytic rates associated with canonical CBMs.

Conclusions

In conclusion, *B/Man5_8*, which is conserved in the *B. animalis* subsp. *lactis* displays a different modular organization, product profile and substrate preference than characterized β -mannanases from gut bacteria. *B/Man5_8* is the first characterized GH5 β -mannanase from the gut niche, which moreover possesses a novel low-affinity soluble-mannan-specific CBM10 not previously described in β -mannanases. This CBM may have evolved to increase the available substrate binding surface, while imparting less reduction in the catalytic rate typically observed for canonical moderate affinity CBMs. These unique substrate-binding properties highlight the diversity of mannan utilization strategies. Further studies are required to assess how these differences and the interplay of extracellular enzymes and transport systems contribute to establishing the hierarchy of mannan degradation in the gut niche.

Additional files

Additional file 1: Composition of the polysaccharide substrates used in this study [56]. (PDF 50 kb)

Additional file 2: Primers used for cloning of the full-length (*B/Man5_8*) and generation of the truncated (*B/Man5_8* Δ CBM10) constructs. (PDF 49 kb)

Additional file 3: pH optimum (A) and stability (B) for *B/Man5_8* activity towards LBG. pH stability is measured as relative activity after 4 days of storage in Britton-Robinson buffers of pH 2-10. Grey error bars represent deviation between duplicate samples. (TIF 1113 kb)

Additional file 4: Temperature optimum (A) and Arrhenius plot (B) for *B/Man5_8*-catalyzed hydrolysis of LBG. Error bars represent deviation between duplicate samples. (TIF 1288 kb)

Additional file 5: Baseline-corrected DSC thermogram of *B/Man5_8* unfolding. Unfolding was measured by the normalized molar apparent heat capacity change (C_p) as a function of temperature. (TIF 3356 kb)

Additional file 6: Michaelis-Menten plots of (A) LBG and (B) M_5 kinetics. The initial normalized rate (V/E_0) data for *B/Man5_8* and *B/Man5_8* Δ CBM10 are depicted as circles and triangles, respectively. The solid lines and the dashed lines depict the fits of the Michaelis-Menten equation to the initial normalized rate data for *B/Man5_8* and *B/Man5_8* Δ CBM10, respectively. (TIF 82 kb)

Additional file 7: Affinity gel electrophoresis with LBG and HEC. In all gels, BSA is in the left lane, *B/Man5_8* in the middle lane, and *B/Man5_8* Δ CBM10 in the right lane. (TIF 1882 kb)

Additional file 8: Relative retardation of *B/Man5_8* in LBG-IV affinity electrophoresis. Relative retardation is measured compared to BSA. The curve fitting was done with nonlinear regression using Prism 6 (GraphPad Software, La Jolla, CA, USA). (TIF 32 kb)

Additional file 9: Sequence alignment of GH5 mannanases. Multiple sequence alignment between characterized β -mannanases representative of different GH5 subfamilies, performed using MAFFT 7 and rendered with ESPrnt 3. The seven residues which are strictly conserved in all of GH5 [39] are marked with green boxes. Trp¹⁹⁶, Trp²⁰⁰ and Tyr²²⁵, the three aromatic residues that are present in the putative aglycone subsites of *B/Man5_8*, are also indicated. (TIF 29861 kb)

Additional file 10: Multiple sequence alignment of GH5_8 sequences. Included are sequences listed in CAZy as having UniProt entries. Positions corresponding to Trp¹⁹⁶ and Trp²⁰⁰ in *B/Man5_8* (UniProt ID: C6A9J9) are indicated. The alignment was done with MAFFT 7 and rendered with ESPrnt 3. (TIF 17898 kb)

Additional file 11: Comparison of kinetic parameters on LBG from characterized β -mannanases. k_{cat} values have been re-calculated as s^{-1} rather than $\text{U} \cdot \text{mg}^{-1}$ where relevant [57–59]. (PDF 54 kb)

Additional file 12: CBM10s of putative and characterized enzymes listed in dbCAN. The list was manually curated by removing three CBM10 sequences with E-values above 0.02 and two which appeared fragmented, and adding the *S/Man5A* (GenBank ID: AAA26710.2) CBM10 sequence. Characterized enzymes are indicated in bold. Enzymes with at least one CBM10 belonging to the same clade as the *B/Man5_8* CBM10 (GenBank ID: ACS46797.1) (Fig. 6) are highlighted with light grey background. (PDF 103 kb)

Additional file 13: Multiple sequence alignment of characterized and predicted CBM10 sequences. The list of CBM10 sequences was obtained from the annotation software website dbCAN (<http://csbl.bmb.uga.edu/dbCAN/index.php>) with the exception of the *S/Man5A* CBM10 sequence which was added manually. The alignment was done with MAFFT 7 and rendered with ESPrnt 3. The positions corresponding to Cys⁵, Tyr⁸, Trp²², Trp²⁴ and Cys³⁶ in the *C/Xyn10A* CBM10 structure [42] are indicated. The four cysteine residues form two disulphide bridges, while Tyr⁸, Trp²² and Trp²⁴ where shown to mediate binding to insoluble cellulose in the *C/Xyn10A* CBM10. (TIF 20792 kb)

Abbreviations

Avicel: Microcrystalline cellulose; CAZy: Carbohydrate-Active enZYmes database; CBM: Carbohydrate-binding module; GG: Guar gum; GH: Glycoside hydrolase; HEC: Hydroxyethyl cellulose; HPAEC-PAD: High-performance anion exchange chromatography with pulsed amperometric detection; INM: Ivory nut mannan; KGM: Konjac glucomannan; LBG: Locust bean (carob) galactomannan; LBG-IV: Low-viscosity locust bean galactomannan; M_1 : Mannose; M_2 : Mannobiose; M_3 : Mannotriose; M_4 : Mannotetraose; M_5 : Mannopentaose; M_6 : Mannohexaose; M_7 : Mannoheptaose; M_8 : Mannooctaose; RU: Response units; SPR: Surface plasmon resonance.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

S.L. provided genomic DNA for *Bifidobacterium animalis* subsp. *lactis* BI-04. J.M., E.K., A.M.S. and M.A.H. performed the experiments. J.M., E.K., H.S., B.S. and M.A.H. planned experiments, interpreted the data and wrote the paper. All authors have read and approved the final version of the paper.

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